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LALIGN finds the best local alignments between two sequences version 2.0u66 September 1998 Please cite: X. Huang and W. Miller (1991) Adv. Appl. Math. 12:373-381

```
Comparison of:
(A) ./wwwtmp/lalign/.18855.1.seq SEQ ID 2
(B) ./wwwtmp/lalign/.18855.2.seq SEQ ID 14
 using matrix file: BL50, gap penalties: -14/-4
  61.1% identity in 108 aa overlap; score:
                                        456 E(10,000): 7.4e-34
        10
                 20
                           30
                                    40
                                             50
                                                      60
      LHGSGSGNEVIEGPQNATVLKGSQARFNCTVSQGWKLIMWALSDMVVLSVRPMEPIITND
SEQ
      SEQ
      {	t LTASGSSYQIIEGPQNVTVLKDSEAHFNCTVTHGWKLLMWTLNQMVVLSLTTQGPIITNN}
       20
                30
                                   50
                                            60
                                                     70
        70
                 80
                           90
                                   100
                                            110
      RFTSQRYDQGGNFTSEMIIHNVEPSDSGNIRCSLQNSRLHGSAYLTVQ
SEQ
                SEO
      RFTYASYNSTDSFISELIIHDVQPSDSGSVQCSLQNSHGFGSAFLSVQ
                90
                         100
                                  110
                                           120
 53.3% identity in 92 aa overlap; score:
                                       290 E(10,000): 4.5e-19
          300
                      310
                               320
                                        330
                                                  340
                                                           350
SEQ
      KKSEKEKTNK---ETETESGNENSGYNSDEQKTTDTASLPPKSCESSDPEQRNSSCGPPH
                  *** **** ***** ********* * * * *****
      .:: . .:::
SEQ
      RKSANMRTNKADPETKLKSGKENYGYSSDEAKAAQTASLPPKSAEVSLPEKRSSSL--PY
          140
                   150
                            160
                                               180
                                                          190
             360
                      370
                               380
SEQ
      QRADQRPPRPASHPQASFNLASPEKVSNTTVV
        SEQ
      QELNKHQPGPATHPRVSFDIASPOKVRNVTLV
            200
                     210
                              220
```

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## | lalign output for SEQ ID 4 vs. SEQ ID 14

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```
Comparison of:
(A) ./wwwtmp/lalign/.19190.1.seq SEQ ID 4
(B) ./wwwtmp/lalign/.19190.2.seq SEQ ID 14
 using matrix file: BL50, gap penalties: -14/-4
  61.0% identity in 105 aa overlap; score: 447 E(10,000): 4.8e-33
           20
                    30
                             40
                                      50
      {\tt SGSGNEVIEGPQNARVLKGSQARFNCTVSQGWKLIMWALSDMVVLSVRPMEPIITNDRFT}
SEQ
      SEQ
      SGSSYQIIEGPQNVTVLKDSEAHFNCTVTHGWKLLMWTLNQMVVLSLTTQGPIITNNRFT
             30
                                                  70
           80
                    90
                            100
                                     110
      SQRYDQGGNFTSEMIIHNVEPSDSGNIRCSLQNSRLHGSAYLTVQ
SEQ
             SEQ
      YASYNSTDSFISELIIHDVQPSDSGSVQCSLQNSHGFGSAFLSVQ
             90
                      100
                               110
  53.3% identity in 92 aa overlap; score:
                                       290 E(10,000): 4.5e-19
      300
                  310
                           320
                                    330
                                             340
                                                       350
SEO
      KKSEKEKTNK---ETETESGNENSGYNSDEQKTTDTASLPPKSCESSDPEQRNSSCGPPH
                  .:: . .:::
      {\tt RKSANMRTNKADPETKLKSGKENYGYSSDEAKAAQTASLPPKSAEVSLPEKRSSSL--PY}
SEQ
          140
                   150
                            160
                                     170
                                                         190
         360
                  370
                           380
SEQ
      QRADQRPPRPASHPQASFNLASPEKVSNTTVV
      :. ... : ::.::..::..:::.:: :.::
SEQ
      QELNKHQPGPATHPRVSFDIASPQKVRNVTLV
           200
                     210
                              220
```

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```
Comparison of:
(A) ./wwwtmp/lalign/.19757.1.seq SEQ ID 6
(B) ./wwwtmp/lalign/.19757.2.seq SEQ ID 14
 using matrix file: BL50, gap penalties: -14/-4
  57.9% identity in 114 aa overlap; score:
                                      458 E(10,000):
          10
                  20
                           30
                                    40
                                             50
      LLTVPEAVGSGSGNEVIEGPQNATVLKGSQARFNCTVSQGWKLIMWALSDMVVLSVRPME
SEQ
      SEO
      LVILAQLTASGSSYQIIEGPQNVTVLKDSEAHFNCTVTHGWKLLMWTLNOMVVLSLTTOG
            20
                     30
                                               60
                                                        70
          70
                  80
                           90
                                   100
                                            110
SEQ
      PIITNDRFTSQRYDQGGNFTSEMIIHNVEPSDSGNIRCSLQNSRLHGSAYLTVQ
                     PIITNNRFTYASYNSTDSFISELIIHDVQPSDSGSVQCSLQNSHGFGSAFLSVQ
SEQ
            80
                     90
                             100
                                     110
                                              120
 53.3% identity in 92 aa overlap; score:
                                     292 E(10,000):
      300
                 310
                          320
                                   330
                                            340
SEQ
      KKSEKEKTNK---ETETESGNENSGYNSDEQKTTETASLPPKSCESSDPEQRNSSCGPPH
                 SEQ
      RKSANMRTNKADPETKLKSGKENYGYSSDEAKAAQTASLPPKSAEVSLPEKRSSSL--PY
         140
                  150
                           160
                                    170
                                                       190
        360
                 370
                          380
SEQ
      QRADQRPPRPASHPQASFNLASPEKVSNTTVV
      SEQ
      QELNKHQPGPATHPRVSFDIASPQKVRNVTLV
           200
                    210
                             220
```

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